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Title:

Allopatric divergence and hybridization within *Cupressus chengiana* (Cupressaceae), a threatened conifer in the northern Hengduan Mountains of western China

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Abstract

Having a comprehensive understanding of population structure, genetic differentiation and demographic history is important for conservation and management of threatened species. High-throughput sequencing (HTS) provides exciting opportunities to address a wide range of factors for conservation genetics. Here, we generated HTS data and identified 266,884 high-quality SNPs from 82 individuals, to assess population genomics of *Cupressus chengiana* across its full range, comprising the Daduhe River (DDH), Minjiang River (MJR) and Bailongjiang River (BLJ) catchments in western China. [Each of ADMIXTURE](#), PCA and phylogenetic analyses indicated that each region contains a distinct lineage, with high levels of differentiation between them (DDH, MJR and BLJ lineages). MJR was newly distinguished compared to previous surveys, and evidence including coalescent simulations supported a hybrid origin of MJR during the Quaternary. Each of these three lineages should be recognized as an evolutionarily significant unit (ESU), due to isolation, differing genetic adaptations and different demographic history. Currently, each ESU faces distinct threats, and will require different conservation strategies. Our work shows that population genomic approaches using HTS can reconstruct the complex evolutionary history of threatened species in mountainous regions, and hence inform conservation efforts, and contribute to the understanding of high biodiversity in mountains.

Keywords: Population genomics, Mountainous regions, Threatened species, ESUs, Hybridization

1 **Introduction**

2 Resolving taxonomic uncertainties and identifying conservation units (CUs) are crucial for the
3 conservation of biological diversity, providing managers and policy makers with a clear
4 understanding of the population unit boundaries of endangered species (Funk, McKay,
5 Hohenlohe, & Allendorf, 2012). Accurate determination of taxonomic status can avoid both
6 underestimation of the necessary protection status for endangered species, and wasted effort on
7 abundant species (Frankham, Ballou, & Briscoe, 2010). Moreover, a species may comprise
8 several genetically distinct evolutionary units, each of which warrants conservation in its own
9 right (Palsbøll, Berube, & Allendorf, 2007). In recent decades, genetic markers that have been
10 used to define CUs have included microsatellite loci (SSR) (Wang, Liang, Hao, Chen, & Liu,
11 2018a), nuclear DNA (nrDNA) (Shang et al., 2015), chloroplast DNA (cpDNA) (Feng, Xu, &
12 Wang, 2018; Petit, El Mousadik, & Pons, 1998) and mitochondrial DNA (mtDNA) (Moritz,
13 1994; Torres-Cambas, Ferreira, Cordero-Rivera, & Lorenzo-Carballa, 2017). However, these
14 markers only yield a few variable loci, and so are generally inadequate for characterizing the
15 population genetic structure of species with complex demographic history and adaptive patterns
16 (Funk et al., 2012).

17 Hybridization, including subsequent introgression, either between species or across
18 intraspecific lineages, can complicate the identification of taxonomic and conservation units,
19 and hence the assignment of priorities when allocating conservation efforts (Allendorf, Leary,
20 Spruell, & Wenburg, 2001; Naciri & Linder, 2015). Nevertheless, hybridization among
21 diverging lineages is prevalent in nature, and about 25% of plant and 10% of animal species are
22 known to have undergone hybridization (Mallet, 2007). Hybridization becomes a conservation
23 issue when gene flow erodes population distinctions, especially when the distinctness of a rare
24 species or race is threatened by introgression from a commoner, sometimes alien, species or
25 race (Allendorf et al., 2001). Equally, however, hybridization is increasingly recognized as a
26 generator of adaptation and biodiversity (Lamichhaney et al., 2018; Rieseberg, 2019). For
27 example, adaptive traits transferred between species by introgression can promote adaptive
28 radiations (Edelman et al., 2019; Rieseberg, 2019). Combining distinct genomes in novel ways,
29 coupled with stabilization and isolation from parents, may form new lineages or species very
30 quickly (Barrera-Guzman, Aleixo, Shawkey, & Weir, 2018; Lamichhaney et al., 2018), which
31 can result in rapid speciation. Thus, hybridization has played an important role in the evolution
32 of many species/lineages (Goulet, Roda, & Hopkins, 2017). Therefore, policy making requires
33 an understanding of the roles hybridization has played in any threatened species. However,
34 detection of hybridization is difficult using traditional molecular markers (Allendorf,
35 Hohenlohe, & Luikart, 2010), with hundreds of markers usually required for accurate
36 determination of the dynamics of hybridization (Allendorf, Hohenlohe, & Luikart, 2010).
37 Recently, the application of high-throughput sequencing (HTS) technologies has made rapid
38 collection of genomic data much easier (Funk et al., 2012), providing exciting opportunities to
39 quantify adaptive variation (Hämälä & Savolainen, 2019; Ma et al., 2019), accurately delimit
40 taxa within critical species complexes (Fennessy et al., 2016; Liu et al., 2018) and assess
41 complex genetic structure, including the effects of hybridization (Ru et al., 2018; Sun et al.,
42 2018; vonHoldt, Brzeski, Wilcove, & Rutledge, 2018). This expanded genomic data will permit
43 many new questions to be addressed regarding conservation (Allendorf, Hohenlohe, & Luikart,

2010), which will make the conservation and management of threatened species more effective.

The Hengduan Mountains (HDMs) region, at the eastern edge of the Qinghai-Tibetan Plateau (QTP), possesses exceptional richness in plant diversity, with about 12,000 species in 1500 genera of vascular plants (Li & Li, 1993; Liu, Duan, Hao, Ge, & Sun, 2014b; Wu, 1988), of which >3300 species (>27.5%) and 90 genera (>6%) are endemic (Sun, Zhang, Deng, & Boufford, 2017). Many of these occur in specific habitats that are also rare and threatened, e.g. *Larix mastersiana*, *Cephalotaxus lanceolata* and *Parakmeria omeiensis* (Fu, 1992; Yong, Bing, & Njenga, 2017). The genetic structure and demographic history of species in the HDMs have been shaped by local orogenetic events and climate oscillations (Favre et al., 2015; Liu et al., 2014b). Mountain uplifts generated geographic barriers that limited gene flow among populations, affecting divergence of lineages, genetic structure, and the evolution of alpine plants (Liu, Sun, Ge, Gao, & Qiu, 2012; Shahzad, Jia, Chen, Zeb, & Li, 2017; Wen, Zhang, Nie, Zhong, & Sun, 2014). This region was also affected by a series of Quaternary glaciations (Zheng, Xu, & Shen, 2002; Zhou & Li, 1998), among which the two largest on the QTP were the Xixiabangma Glaciation and the Naynayxungla Glaciation, which occurred around 1.2-0.8 million years ago (Mya) and 0.72-0.5 Mya, respectively (Zheng et al., 2002; Zhou & Li, 1998). Many tree species on the QTP moved south and/or to lower altitudes during the ice ages (Liu et al., 2014b; Qiu, Fu, & Comes, 2011), which could drive intraspecific divergence, or hybridization if diverged lineages share a refugium (Du, Hou, Wang, Mao, & Hampe, 2017; Liu, Abbott, Lu, Tian, & Liu, 2014a; Liu et al., 2013; Ren et al., 2017; Sun et al., 2014). Hence species distributed in the HDMs may have complex evolutionary histories, necessitating large numbers of markers to accurately delimit both closely related species and intraspecific lineages.

The Minjiang Cypress, *Cupressus chengiana* S.Y. Hu, is a threatened conifer that occurs around the northern HDMs, where it is a vital ecological component of arid valley ecosystems, and is regularly used for house construction and furniture production. It has suffered a sharp decline in range and population size because of logging (Hao et al., 2006; Zeng & Yang, 1992), and is now classified as “Vulnerable” by the IUCN (Zhang & Christian, 2013), and as a “Second-Class Endangered Plant” of China (Fu, 1992). Early studies using three regions of chloroplast genome (Xu et al., 2010), six pairs of nuclear microsatellite markers (Lu et al., 2014) and ten nuclear DNA sequence loci (Xu et al., 2017) demonstrated clear genetic differentiation between Bailongjiang river material in Gansu province (hereafter labelled BLJ) and material from the Daduhe and Minjiang rivers in Sichuan Province. This suggested that *C. chengiana*, comprises two evolutionary significant units (ESUs): one in BLJ, and the other Daduhe (DDH) plus Minjiang (MJR). The BLJ material would currently satisfy the IUCN (2012) criterion of “Endangered” if treated alone. However, currently available data cannot provide a comprehensive understanding of its genetic status, and therefore population genomic data are needed to address broader factors of conservation for this threatened species. Here, we collected HTS data to characterise genetic variation across *C. chengiana* populations to address the following questions. (i) How many ESUs can be identified within *C. chengiana* based on HTS data? (ii) What roles have past environmental changes and hybridization played in its evolutionary and population history? (iii) Do adaptive differences exist among the ESUs? (iv) What conservation implications and recommendations can be inferred from our data, for this rare conifer? A robust inference for the genetic status and lineage evolutionary history of *C.*

chengiana would facilitate conservation and management of this threatened species, as well as shedding light on the evolution of species and [populations](#) within the HDM biodiversity hotspot.

Materials and Methods

Sampling and RNA sequencing

Cupressus chengiana is now restricted to three isolated arid valleys between 800 and 2900 m a.s.l. in the upper reaches of the Daduhe (DDH), Minjiang (MJR) and Bailong (BLJ) rivers (Fu, Yu, & Farjon, 1999; Hao et al., 2006; Xu et al., 2017). The Minjiang river material lies roughly between the other two regions in both location and altitude, and is separated from Bailongjiang and Daduhe rivers by the Minshan and Qionglai Mountains, respectively. We collected across the full range of *C. chengiana* from 2016 to 2018, and sampled fresh leaves in thirteen locations across the three river catchments for RNA-seq: 35 individuals for BLJ, 17 for MJR, and 30 for DDH (Table 1, Figure 1). In each location, the distance from every sampled individual to any other was more than 50m, to avoid the impact of potential clonal reproduction. Five samples each of *C. duclouxiana* and *C. gigantea* that were collected in our previous work (Ma et al., 2019) were included as outgroups.

Fresh leaves were put in liquid nitrogen immediately and kept below -80°C before extraction. RNeasy Pure Plant Plus Kits (TIANGEN® Biotech, Beijing, China), which provide an efficient method for purification of total RNA from plant tissues rich in polysaccharides and polyphenolics, was used to isolate total RNAs. A NanoPhotometer® spectrophotometer (IMPLEN, CA, USA) was used to check RNA purity, and a Qubit® RNA Assay Kit in Qubit® 2.0 Fluorometer (Life Technologies, CA, USA) was used to measure RNA concentration. RNA integrity was assessed via the RNA Nano 6000 Assay Kit of the Agilent Bioanalyzer 2100 system (Agilent Technologies, CA, USA).

Sequencing libraries were generated using a NEB Next® Ultra™ RNA Library Prep Kit for Illumina® (NEB, USA) following manufacturer's recommendations. Briefly, index codes were first added to attribute sequences to each sample, and then mRNAs were fragmented into short sequences. cDNA was synthesized for each RNA [fragment](#), and NEBNext Adaptors with hairpin loop structure were ligated to prepare for each cDNA fragments. PCR was then performed with Phusion High Fidelity DNA polymerase. Finally, PCR products were purified and library quality was assessed on the Agilent Bioanalyzer 2100 system. After the above steps, the library preparations were sequenced on Illumina HiSeq X Ten platforms to generate 150bp paired-end raw reads.

SMRT sequencing of full-length transcriptome

To get a high-quality reference, single-molecule real-time (SMRT) sequencing was used to obtain the full-length transcriptome of *C. chengiana*. Fresh leaves, stems and female seed cones of one individual were sampled and then snap-frozen in liquid nitrogen. After RNA extraction and quality checking (see above), pooled RNA comprising equal amounts of high-quality RNA from the three tissues was used for cDNA synthesis and library construction using a SMART PCR cDNA kit (Clontech, Mountain View, CA, USA) and the BluePippin Size Selection System. This library was subsequently sequenced on a Pacific Biosciences (PacBio) RS sequencing instrument.

Raw reads of PacBio full-length isoform sequencing were processed using the SMRT Link ver. 4.0 software (<https://www.pacb.com/support/softwaredownloads>). From this, Circular Consensus Sequences were generated; these were then classified into full length and non-full length reads by examining for poly(A) signal and 5' and 3' adaptors. Consensus isoforms were identified from full-length non chimera sequences (FLNC), and polished with non-full length reads to obtain high-quality isoforms (HQ, above 99% accuracy) using the Quiver algorithm from SMRT Link. The Illumina RNA-seq data from the same individual was used to correct the PacBio sequences performed in LoRDEC (Rivals & Salmela, 2014).

To eliminate confounding effects from microbial and plastid DNA, we removed sequences showing high similarity with either microbial DNA sequences (MBGD, downloaded from http://mbgd.genome.ad.jp/htbin/view_arch.cgi (Uchiyama, Higuchi, & Kawai, 2010) or any part of the complete chloroplast genome of *C. jiangeensis* (GenBank accession: NC_036939.1) (Li et al., 2019). The HQ full-length polished consensus transcripts had their redundancy removed by CD-HIT-EST ver. 4.6.1 (Li & Godzik, 2006), and were then processed with Cogent ver. 3.1 (<https://github.com/Magdoll/Cogent>) to obtain a final set of unique transcript isoforms (referred to as UniIsoforms).

Read mapping and SNP calling

Illumina raw reads were filtered via Trimmomatic ver. 0.36 (Bolger, Lohse, & Usadel, 2014). This involved first removing adapters or bases from either the start or the end of reads with base Phred quality score (Q) < 3, and then discarding poly-N reads (those with >10% unidentified nucleotides) and low-quality reads (those with over 50% of bases with Q < 3). Finally, reads with more than 36 bases after trimming were retained as quality-filtered reads.

We used BWA-MEM ver. 0.7.12 (Li & Durbin, 2009) with default parameters to align the quality-filtered reads of each individual to the nuclear transcriptome sequences (UniIsoforms). SAMTOOLS ver. 1.2 (Li et al., 2009a) was run to convert Sequence Alignment/Map (SAM) files to Binary Alignment/Map (BAM) files, and sort BAM files. We used PICARDTOOLS ver. 2.8.1 (<http://broadinstitute.github.io/picard/>; Broad Institute, GitHub Repository) to mark and remove duplicate reads. The regions around indels were realigned using the RealignerTargetCreator and IndelRealigner tools in GATK ver. 3.7 (DePristo et al., 2011). We used the “mpileup” command in SAMTOOLS (Li et al. 2009) to identify SNPs with parameters “-q 20 -Q 20 -t AD,ADF,ADR,DP,SP”. Data were filtered with the following processes: SNPs with a mapping quality <30, a mapping depth <10, genotyping rate <50% per group, minor allele frequency (MAF) <5%, or in 5bps windows around any indel. The program SnpEff (Cingolani et al., 2012) was used to annotate SNPs.

Genetic structure and phylogenetic inference

We used VCFtools (Danecek et al., 2011) and a perl script (Ru et al., 2018) to estimate the value of Tajima's *D*, population genetic differentiation (F_{ST}), absolute differentiation (D_{XY}) and nucleotide diversity (π , for all callable sites). To keep rare variants, the MAF control was not performed for the data set that was used to calculate Tajima's *D* and π .

A model-based evolutionary clustering analysis via ADMIXTURE ver. 1.23 (Alexander &

Lange, 2011) was used to identify evolutionary clusters. We used VCFtools and PLINK ver. 1.90 (Purcell et al., 2007) to convert input data and remove linkage disequilibrium sites with the parameter set as “--indep-pairwise 50 5 0.4”. The most likely number of genetic clusters (K) was estimated in ADMIXTURE ver. 1.23, by computing parameters’ maximum-likelihood estimates. Ten independent simulations were run for each value of K from one to ten with cross validation to investigate the convergence of samples. The minimization of cross-validation error among all runs was used to determine the most likely number of clusters. In order to compare with results from ADMIXTURE, principal component analysis (PCA) on *C. chengiana* individuals was conducted to explore the species’ genetic structure, using the SMARTPCA program in the software EIGENSOFT ver. 6.1.3 (Price et al., 2006).

A perl script (Ru et al., 2018) was then used to generate concatenated sequences of each individual. Here, only neutral sites (4DTv, four-fold degenerate sites) were retained to construct phylogenetic inference. The software jModelTest (Darriba, Taboada, Doallo, & Posada, 2012) was used to select the best-fit model of nucleotide substitution using Akaike Information Criterion. Maximum-likelihood (ML) trees were reconstructed in RAxML ver. 8.2.9 (Stamatakis, 2014) using *C. duclouxiana* and *C. gigantea* as outgroups. We performed 200 bootstrap replicates to calculate the node support values.

Phylogenetic-network analysis

To obtain single-copy genes, one individual from each of the three groups (BLJ, MJR and DDH) within *C. chengiana*, plus one *C. gigantea* accession, were selected for orthologous sequences searching. Quality-filtered reads were assembled into contigs in Trinity ver. 2.8.4 (Grabherr et al., 2011) with default parameters. We used the BUSCO (Simão, Waterhouse, Ioannidis, Kriventseva, & Zdobnov, 2015) database to assess the transcriptome assembly. The longest transcript for each gene was selected by a custom python script (see Supplemental Information), and then we used CD-HIT-EST ver. 4.6.1 (Li & Godzik, 2006) to eliminate redundancies. Coding and peptide sequences were predicted by TransDecoder ver. 5.5.0 (Haas et al., 2013). The 1:1:1:1 orthologous gene data set was generated for BLJ, MJR, DDH and *C. gigantea* (outgroup) in Orthofinder ver. 2.3.3 (Emms & Kelly, 2015). The corresponding coding sequences of each orthogroup were aligned via MAFFT (Katoh & Standley, 2013), and trimAL ver. 1.4.1 (Capella-Gutiérrez, Silla-Martínez, & Gabaldón, 2009) was used to remove positions with more than 50% missing data. Those aligned sequences that were longer than 300 nucleotides were retained to generate the best rooted ML trees in RAxML under the GTR+GAMMA substitution model using rapid-bootstrapping approach. Meanwhile, we used ASTRAL ver. 5.6.3 (Mirarab et al., 2014) to estimate the species tree with 100 bootstrap replicates. Finally, a set of 10,227 orthologous gene trees was examined using PhyloNet ver. 3.6.10 (Than, Ruths, & Nakhleh, 2008) to infer reticulate evolutionary relationships for *C. chengiana*. A custom python script (see Supplemental Information) was used to convert the format for input files. We used the command InferNetwork_ML_Bootstrap with the parameter “InferNetwork_ML_Bootstrap 2 -pl 6 -di” to infer a species network, where the maximum number of reticulations was set as 2 and the sampling process was repeated 100 times in parametric bootstrap by default.

Demographic modelling and gene flow

Although some synonymous sites are expected to evolve under purifying selection (Lawrie, Messer, Hershberg, & Petrov, 2013), they are generally assumed to be under weak selection and nearly neutral (Akashi, 1995; Yang & Nielsen, 2008). Therefore, many studies used 4DTv sites to minimize the bias in demographic inferences when more neutral sites were unavailable (Marburger et al., 2019; Zhang et al., 2017). Here, we also used SNPs at 4DTv sites for demographic inference to reduce the impact of natural selection. SNPs without MAF filtering were further filtered to remove all missing data across all individuals sampled. We used a perl script (Ru et al., 2018) to generate folded two-dimensional joint site frequency spectra (2D-SFS). The 2D-SFS for all *C. chengiana* individuals was estimated by *fastsimcoal2* (FSC2) (Excoffier, Dupanloup, Huerta-Sánchez, Sousa, & Foll, 2013). We used the Akaike information criterion (AIC) to rank all models, and chose the model that was the best fit to the data, to reduce subjective bias. Based on early results confirming that MJR was roughly intermediate between the other two populations, we selected four possible scenarios, and compared 11 possible models based on these. The scenarios were: (i) a hybrid origin of the MJR lineage from the other two lineages (models 1-3, Figure S1); (ii) divergence of the MJR and BLJ lineages from a recent common ancestor (((MJR, BLJ), DDH), models 4-6, Figure S1); (iii) divergence of the MJR and DDH lineages from a recent common ancestor (((MJR, DDH), BLJ), models 7-9, Figure S1); and (iv) radiative evolution among all three lineages ((MJR, DDH, BLJ), models 10-11, Figure S1). The mutation rate was set as 9.7×10^{-9} per site per generation following Li et al. (2012)'s estimate for Cupressaceae species. Because of long generation times in gymnosperms (De La Torre, Li, Van de Peer, & Ingvarsson, 2017), we assumed an average generation time of 50 years for *C. chengiana*, which is about three to five times the age at first reproduction but less than the maximum expected lifespan of conifers (Bouillé & Bousquet, 2005). This generation time was also commonly adopted in many studies for conifers (Bouillé & Bousquet, 2005; Li et al., 2009b; Ma et al., 2019). A parameter bootstrapping approach was used to construct 95% confidence intervals (CI) with 50 independent runs. We used the Stairway plot method (Liu & Fu, 2015) to investigate the detailed population demographic history for each lineage using the folded one-dimensional SFS (Figure S2) from 4DTv sequences.

We further used the ABBA-BABA test (*D*-statistics) (Durand, Patterson, Reich, & Slatkin, 2011) to test for the possibility of gene flow among the three groups. Based on the patterns of ancestral and derived alleles in the ingroups and outgroups, this analysis can distinguish between incomplete lineage sorting and hybridization (Elgvin et al., 2017; Zhang et al., 2019). Two topologies ((BLJ, MJR), DDH) and ((DDH, MJR), BLJ) were selected to calculate a *D* value using ANGSD's ABBA-BABA multipopulation tool (Korneliussen, Albrechtsen, & Nielsen, 2014) with five *C. duclouxiana* individuals as the outgroup. We used a Z-test to determine if the *D* value was significantly deviated from zero. We considered Z scores >3 to be significant.

Detection of candidate genes and GO annotation

In endangered species, adaptive differences between populations can create differing ecological requirements, necessitating distinct management strategies for each of them (Crandall, Bininda-Emonds, Mace, & Wayne, 2000). At first, we assumed no *a priori* information in order to examine adaptive differentiation patterns for *C. chengiana*, and to test if there were genes under strong directional selection in any set of populations (Funk, McKay, Hohenlohe, & Allendorf,

2012). We performed a global F_{ST} outliers test in BAYESCAN ver. 2.1 (Foll & Gaggiotti, 2008) with default parameters. A subpopulation specific fixation index (F_{ST}) was used to estimate the difference in allele frequency between the total population (all individuals of *C. chengiana*) and each subpopulation (BLJ, MJR and DDH). To reduce the false discovery rate when making decisions, q-values were calculated in BAYESCAN. Among outliers, those having a q-value lower than 0.001 were treated as false positives, and hence the remaining 99.9% of corresponding outliers were expected to not be false positives.

DDH group occupies the highest habitats (>2000m) among the three groups of *C. chengiana*. Therefore, we further used the population branch statistic (PBS) (Yi et al., 2010) to identify candidate genes related to high-altitude adaptation in DDH, comparing to the lowland BLJ group (<1500m). Because individuals of *C. duclouxiana* and *C. gigantea* were clustered together as a clade in an outgroup position to *C. chengiana* (Figure 2c, 2d), all ten individuals of *C. duclouxiana*+*C. gigantea* were treated as a single outgroup population. First, pairwise F_{ST} values were calculated in VCFtools, and the population divergence time T , in units scaled by the population size, was obtained as $T = -\log(1-F_{ST})$ (Cavalli-Sforza 1969). Next, the length of the branch leading to the DDH population since the divergence from the BLJ was estimated as:

$$PBS_{DDH} = \frac{T^{DDH-BLJ} + T^{DDH-outgroup} - T^{BLJ-outgroup}}{2}.$$

Genes with the highest 1% of PBS were recognized as highly divergent genes, which could result from positive selection (Wang et al., 2018b).

The software ANGEL ver. 2.4 (<https://github.com/PacificBiosciences/ANGEL>) was used to predict open reading frames, and translate protein codes. We used BLASTP ver. 2.2.23 (Altschul et al., 1997) to compare the protein sequences, using the Swiss-Prot protein sequence database for homology search analysis. Gene ontology (GO) terms for each gene were searched using the Blast2GO program (Conesa et al., 2005). For the functions of genes and gene set enrichment analysis, the analysis tool Singular Enrichment Analysis (SEA) in agriGO ver. 2.0 (Tian et al., 2017) was used. The Chi-squared test was used to calculate the statistical significance of enrichment, with P -values below than 0.05 treated as significant following adjustment via the Benjamini–Yekutieli procedure to control for false discovery rate.

Chloroplast phylogenetic analysis

Previous research has shown that chloroplast (cp) sequences represent a large fraction of the plant transcriptome (Osuna-Mascaró, Rubio de Casas, & Perfectti, 2018; Ru et al., 2018). We used the BWA-MEM algorithm of BWA ver. 0.7.12 (Li & Durbin, 2009) to map quality-filtered reads of each individual against the published complete cp genome sequence of *C. jiangensis* (Li et al., 2019) to examine cpDNA variation. After removing duplicate reads, and realigning regions around indels (see above), SAMTOOLS (Li et al. 2009) was used to identify SNPs. SNPs were filtered with the following processes: SNPs were removed if they had a mapping quality <30, a mapping depth <3, genotyping rate <50% per group, minor allele frequency (MAF) <5%, or in 5bps windows around any indel. Concatenated sequences of each individual were used to reconstruct ML trees in RAxML using *C. duclouxiana* and *C. gigantea* as outgroups.

Ecological niche modelling

Current potential distributions for each group were predicted using ecological niche modelling (ENM) in MAXENT ver. 3.3.4 (Phillips & Dudík, 2008) with the parameters set as “replicates: 20 replicates; type: subsample; maximum iterations: 5000; random test points: 25”. Climate layers comprising 19 bioclimatic variables of a 2.5 arc minute resolution were downloaded from WorldClim database (version 1.4, <http://www.worldclim.org>), and in addition, one altitude layer was downloaded from the SRTM elevation database (<https://www2.jpl.nasa.gov/srtm/>). We calculated pairwise Pearson’s correlation coefficients (r) (Dormann et al., 2013) for current climate and altitude data across distributions of all trees performed in ENMTools ver. 1.4.3 (Warren, Glor, & Turelli, 2008; 2010). Any factor that had a correlation coefficient greater than 0.7 with two or more other factors was excluded. The geographic coordinates of 18 locations from DDH, 13 from MJR and 14 from BLJ (Table S1) were collected from field investigations or previous publications (Xu et al., 2010, 2017), which covered most of the known area of occupancy of this species; these were inputted into MAXENT. We conducted a hierarchical partitioning approach (Chevan & Sutherland, 1991) to confirm which variable independently contributed most, using the R package hier.part (Walsh, Mac Nally, & Walsh, 2003). The performance of models was predicted by comparing their AUC values (the area under the receiver operating characteristic curve). AUC values range from 0 to 1, where a score of 1 indicates perfect discrimination (Fielding & Bell, 1997). Niche overlap and identity tests were performed in ENMTools to measure niche differences between groups by calculating Schoener’s D (Schoener, 1968) and standardized Hellinger distance (I). The values of D and I both ranged from 0 to 1, which indicated no niche overlap or identical niches respectively.

To further examine the patterns of distribution shifts within *C. chengiana*, we also used ENM to predict potential distributions during the Last interglacial (LIG, ~120,000 - 140,000 years ago), the last glacial maximum (LGM, about 22,000 years ago) and the future (2050, average for 2041-2060) for each group. For the period of LGM, layers of four models available at the WorldClim database were downloaded to generate average-over-pixel bioclimatic variables following Zheng et al. (2017). Future climate data was available from the Fifth Phase of the Coupled Model Intercomparison Project (CMIP5), while the climate data during LIG was downloaded from WorldClim database (source: Otto-Bliesner et al. 2006).

Results

Full-length transcriptome analysis using PacBio Iso-Seq

Using mixed RNA samples of leaf, stem and female cone, we obtained 19.32G of nucleotide (nt) reads of inserts (ROIs) from three SMRT cells. The number of ROIs was 13,637,084, and the mean length was 1,417nt. The Iso-seq classification and clustering protocol yielded 47,546 polished high-quality (HQ) transcripts, while the N50 was 3,117nt (Table S2). UniIsoforms were excluded from the final set if they had similarity to either microorganisms or the plastid genome that was used as a reference. The total size of the reference UniIsoforms data set was 50.506M nt, and the N50 was 3,133nt (Table S2).

SNP calling

After removing low quality sequences, 3.4 billion filtered-quality reads were obtained for the

82 individuals from an Illumina platform. By mapping these filtered-quality reads to the reference UniIsoforms, we identified 5.82 million nuclear SNPs. After quality control, a total of 266,884 high-quality nuclear SNPs was retained. A total of 5,202 cpDNA SNPs was successfully identified using the complete chloroplast genome of *C. jiangensis* as a reference. After all filtering steps, we finally retained 1,251 SNPs from which to reconstruct cp phylogenetic trees.

Population genetic structure and genetic diversity

Three distinct genetic clusters were detected by both PCA and ADMIXTURE analyses. From the PCA plot, the first principal component (PC1), which explained 12.26% of all genetic variance, differentiated the three geographically distinct *C. chengiana* groups: MJR, BLJ and DDH, with MJR occupying an intermediate space between BLJ and DDH according to PC1 (Figure 2b). Results of ADMIXTURE also indicated that three genetic groups ($K=3$) were optimal (Figure S3). For $K=3$, a clear genetic differentiation among the same three groups was detected, with the clearest differentiation between DDH and MJR (Figure 2a). In the scenario of $K=2$, individuals of the BLJ and DDH clades clustered into two distinct groups, while the MJR group contained a mixture of genetic components of the other two groups (Figure 2a), which is consistent with MJR being of hybrid origin.

Of the entire set of 266,884 nuclear SNPs, 11,913 were specific to DDH, 11,489 to BLJ and 3,213 to MJR (Table 2). DDH and BLJ shared the fewest SNPs (14,748), whereas MJR shared more with each of DDH (20,245) and BLJ (27,524, Table 2), consistent with a hypothesis of a hybrid origin for MJR.

Regarding population differentiation, genetic distance was highest between BLJ and DDH ($F_{ST}=0.1752$), whereas MJR had F_{ST} values of 0.1066 and 0.1397 with BLJ and DDH, respectively. The average value of absolute divergence (D_{XY}) between BLJ and DDH ($D_{XY}=0.3440$) was also greater than that between MJR and either BLJ ($D_{XY}=0.3008$) or DDH ($D_{XY}=0.3104$, Table 2).

SNPs without MAF filtering were used to calculate the π and Tajima's D values. The average π value for BLJ (0.0069, Table 2) is less than that for MJR (0.0072, Table 2), while DDH has the lowest π value (0.0064, Table 2). The average Tajima's D values are -0.1790, 0.0470, and -0.1449 for BLJ, MJR and DDH respectively (Table 2).

Phylogenetic inference for nuclear and chloroplast SNPs

Based on the results of jModeltest (Table S3), we used the GTR+GAMMA model for ML tree reconstruction. From the phylogeny for nuclear SNPs, three distinct lineages were detected, corresponding exactly to BLJ, DDH and MJR (Figure 2c), with MJR closer to BLJ than to DDH. A coalescent-based species tree generated by ASTRAL produced a very similar result (Figure S4d). In contrast, an ML tree constructed from cp SNPs identified two distinct clades within *C. chengiana*, with one comprising BLJ and the other DDH+MJR (Figure 2d).

Reticulate evolutionary relationships within *C. chengiana*

The contig N50 of the assembled transcriptome for BLJ, MJR, DDH and *C. gigantea* is 1,592, 1,682, 1,595 and 1,143, respectively (Table S4). More than 80% of the genes in the BUSCO

plant set were covered by all four of the assembled transcriptomes (Table S4). A total of 10,233 single copy orthogroups was identified in Orthofinder, and 10,227 of them were retained to reconstruct [gene trees using ML](#). Hence a total of 10,227 gene trees were generated in RAXML, and of these, 3,975 (38.87%) showed the closest relationship between MJR and BLJ (Figure S4a), whereas 3,426 (33.50%) clustered MJR with DDH (Figure S4b), and 2,826 (27.63%) clustered DDH with BLJ (Figure S4c). Results of PhyloNet showed reticulate evolutionary relationships among BLJ, MJR and DDH, indicating a hybrid origin for MJR. The inheritance probability between MJR and BLJ was 58.87%, which was higher than that between MJR and DDH (41.13%) (Figure 3c), indicating a greater genomic contribution of BLJ to MJR.

Demographic history and gene flow

By comparing the AIC values for all 11 models, the hybrid speciation model with continuous migration among the three groups was the best-fitting model (model3, Figure 3b, Table S5). Divergence between BLJ and DDH was dated to (4.23-) 4.56 (-4.87) Mya (incorporating 95% CI; Table 3). The estimated hybrid parameter (α) indicated that ~62% of the nuclear genome of the initial MJR population came from BLJ, and ~38% from DDH, which was consistent with genetic admixture (Table 3; Figure 2a; Table S6). The population sizes for BLJ, MJR and DDH were estimated to be 238,794, 114,433 and 166,952 respectively (Table 3; Table S6). The ancestral effective population size ($N_A=323,866$, Table 3) was estimated to be larger than any of these (Table 3). A stairway plot analysis showed that a decline of population size for DDH occurred from 0.9 to 0.6 Mya, followed by an expansion 0.6-0.4 Mya, coinciding respectively with the Naynayxungla glaciation, and its end (Figure 3a). In contrast, BLJ maintained a stable and high effective population size (~160,000) over the past seven million years (Figure 3a). The population size of MJR expanded rapidly until approximately 10-7 Mya, and declined to ~110,000 around 1.5 Mya, and then maintained that size with little fluctuations.

Asymmetric gene flow between MJR and the other two groups was detected, with the rates of migration from MJR to each of BLJ ($M_{1\leftarrow 2} = 9.22E-6$) and DDH ($M_{3\leftarrow 2} = 1.00E-5$) being higher than those in the opposite direction ($M_{2\leftarrow 1} = 3.65E-6$; $M_{2\leftarrow 3} = 4.94E-6$, respectively) (Figure 3b, Table 3). Results of the ABBA-BABA test suggested that significant gene flow had occurred between MJR and both of BLJ and DDH on the genomic level (Table 4), which was not consistent with the genetic pattern of MJR being the result of incomplete lineage sorting.

Identification and characterization of outlier loci

With a 0.1% threshold for the q-value, we identified 575 outlier SNPs (Figure 3d), which suggested a divergent differentiation, and that these markers could have [been subject](#) to divergent selection among MJR, BLJ and DDH, based on the Bayesian method performed in BAYESCAN. The average F_{ST} estimated in BAYESCAN was (0.1718-) 0.1934 (-0.6522). Nearly 90% of the SNPs (237,792 of 266,844; 89.10%) showed $F_{ST} < 0.2$, while the F_{ST} value for outlier SNPs was high, i.e. (0.4628-) 0.5303 (-0.6522), suggesting that the three groups were indeed greatly differentiated at outlier SNPs. These outliers might putatively be under divergent selection, representing evidence of adaptive [differentiation](#) between the three groups. These outliers were located in 226 genes, of which 157 were annotated in the Swiss-Prot protein sequence database. Gene ontology enrichment analyses of all outliers detected 12 significantly

over-represented GO terms ($P < 0.05$, $FDR < 0.05$), including: “stilbene biosynthetic process”, “coumarin biosynthetic process”, “lignin metabolic process”, “L-phenylalanine metabolic process” and “double-stranded DNA binding” (Table S7).

We further used the PBS approach to identify genes potentially under positive selection in the DDH group. A total of 127 genes (top 1%) were identified in DDH with $PBS_{DDH} \geq 0.7610$, and 74 of them were annotated. In total, 18 significantly over-represented GO terms with corrected P -value < 0.05 were identified (Table S8). Among the 18 GO terms in DDH, six had also been identified by BAYESCAN, including the “stilbene biosynthetic process” and “stilbene metabolic process” (Table S8). Furthermore, although no significant GO category with corrected P -value < 0.05 was found to be involved in the functions of response to abiotic/biotic stresses in highland environments, some genes with extreme PBS_{DDH} exhibited the signature of high-altitude adaptation in DDH. These included six genes involved in “cellular response to DNA damage stimulus”, 11 related to “positive regulation of response to stimulus”, 34 related to “response to abiotic stimulus”, and 14 related to “response to abscisic acid” (Table S9). These genes were also extremely differentiated between DDH and either BLJ or MJR, and different alleles for all of these genes were fixed between DDH and BLJ (Figure S5).

Ecological niche differences among *C. chengiana* populations

ENMs were constructed for the three *C. chengiana* groups to predict their current potential distributions and then the model was projected to past and future scenarios. Seven bioclimatic variables (Alt: altitude, Bio2: mean diurnal range, Bio3: isothermality, Bio4: temperature seasonality, Bio15: precipitation seasonality, Bio16: precipitation of wettest quarter, and Bio19: precipitation of coldest quarter) were retained with $r < 0.7$ in each pair. Values of AUC of all models were 0.989 ± 0.007 for BLJ, 0.984 ± 0.006 for MJR, and 0.998 ± 0.001 for DDH, indicating that all models performed better than random expectation. The environmental variables that showed the highest independent contributions were Bio19 (37.32%), Bio19 (27%), and Alt (22.04%) for BLJ, MJR, and DDH respectively (Figure S6). Observed measures of niche similarity (D and I) were lower than null distributions for DDH vs either BLJ or MJR, suggesting high niche differentiation between DDH and both of BLJ and MJR (Figure 4b). However, D and I fell within the range of null distributions for BLJ vs MJR, suggesting that few niche differences exist between these two (Figure 4b).

For the LIG model, all groupings were predicted to have undergone clear southward range shifts (Figure S7). For both BLJ and MJR individually, northward distribution shifts were predicted for LGM model, while the predicted present-day and LGM distributions were nearly identical for the DDH group (Figure S7). The future model showed clear range expansions for both MJR and DDH relative to the present day, while a clear distribution contraction was predicted for BLJ (Figure S7).

Discussion

Cupressus chengiana comprises three ESUs, and the Minjiang river ESU is of hybrid origin

Here, we employed population genomic data to explore the genetic diversity, genetic structure and demographic history for the threatened conifer *C. chengiana*, to aid in its conservation.

Multiple lines of evidence presented here suggested that material from each of the three river catchments (BLJ, DDH and MJR) forms a distinct genetic lineage, with a high level of genetic differentiation between the three. ADMIXTURE and PCA demonstrated that no overlaps existed between lineages, whereas the ML tree constructed from the nuclear SNPs demonstrated that each lineage was reciprocally monophyletic. These three lineages might represent the early stages of speciation by isolation, and each forms an important component of the conifer diversity of the world. Hence, each of BLJ, DDH and MJR represents an evolutionarily significant unit (ESU).

The major difference from previous results is the clear differentiation between DDH and MJR, which had not been detected in past studies based on limited data (Lu et al., 2014; Xu et al., 2017). Here, our analyses based on population genomic data presented evidence that the newly recognized MJR ESU had a hybrid origin from the other two ESUs. Population genetic analyses indicate that this ESU is genetically admixed between BLJ and DDH (Table 2, Figure 2a), and demonstrate a reticular evolutionary relationship (Figure 3c). Results of coalescent analysis strongly favored a hybrid origin over non-hybrid scenarios (Figures 3b, S1, Table S5). Thus, multiple analyses indicate that the MJR ESU might be a lineage of hybrid origin, with ~62% of its nuclear composition derived from BLJ, and ~38% from DDH (Figures 3b). Detection of admixture signals had been difficult in earlier studies using <10 loci (Allendorf et al., 2010), but HTS data as used here provides abundant markers that can contribute to the accurate description of dynamics of hybridization and introgression (Allendorf et al., 2010; Witherspoon et al., 2007). Our work confirmed the advantages of population genomic approaches using HTS for research concerning conservation genetics.

Demographic history and gene flow among *C. chengiana* lineages

The strong geographic structure here detected for the three *C. chengiana* lineages implied that the Qionglai and Minshan Mountains may have been able to limit gene flow between populations occupying separate valleys, at least in *C. chengiana*. According to the optimal model from FSC2 analysis (Figure 3b, Table 3), divergence between DDH and BLJ occurred (4.23-) 4.46 (-4.87) Mya. This falls within the timescale of the intense uplifts of the HDMs that occurred from the Late Miocene onwards, approaching their highest elevation before the Late Pliocene (Favre et al., 2015; Sun et al., 2011; Xing & Ree, 2017). Other lineage divergence events in the HDMs or QTP during this period, include intraspecific differentiation in *Taxus wallichiana* (~4.2 Mya) (Liu et al., 2013) and *Quercus aquifolioides* (~4.4 Mya) (Du et al., 2017), plus interspecific differentiation between *C. gigantea* and *C. duclouxiana* (~3.35 Mya) (Ma et al., 2019), all of which might be the results of uplift events in this area. If the Qionglai and Minshan Mountains were uplifted at this time, these might have caused the divergence of DDH from BLJ.

The hybridization event that formed the MJR group was estimated to have occurred at (1.14-) 1.34 (-1.41) Mya (Figure 3b), a little before the start of the Xixiabangma Glaciation (around 1.2 Mya). If hybridization occurred during this or the previous glaciation, then it might have been triggered by southward and downhill migration of *C. chengiana* in response to climate cooling. The Daduhe and Minjiang rivers meet at 400 a.s.l., whereas BLJ material might have spread across suitable ground at lower altitudes to meet them. Previous studies showed that the

largest Quaternary glaciation had begun causing alterations to plant distributions in this period (Li & Fang, 1999; Lisiecki & Raymo, 2007; Sun et al., 2014), and the homoploid hybrid species *Picea purpurea* might also have originated during this time (~1.3 Mya) (Sun et al., 2014).

Alternatively, upward migration during a warmer interglacial might have reduced the barrier presented by the Qionglai and Minshan Mountains, and led to contact at higher altitudes. Either way, it is likely that the Minjiang river basin was first occupied by individuals of one lineage (more likely BLJ, based on genetic similarity) and then invaded by genetic material of the other ~1.34 Mya. This is also supported by phylogenomic analyses based on cp genome-wide SNPs, wherein MJR and DDH were clustered together and barely distinguishable, whereas BLJ formed a distinct and separate clade (Figure 2d), in common with Xu et al., (2010). In conifers, cpDNA is transmitted via pollen, and nrDNA biparentally by seeds (Mogensen, 1996; Sun et al., 2014), hence DDH appears to have been the pollen parent of MJR. This fits an idea that BLJ material around the Minjiang river was invaded via DDH pollen to form the MJR lineage. Taken overall, we can infer that both past orogeny and climatic events in HDMs region may have acted as major factors in shaping the evolutionary history of *C. chengiana*.

FSC2 analysis indicates that gene flow between all three populations might have occurred since they diverged (models 2 or 3, Fig S1). Admixture of cpDNA haplotypes between DDH and MJR does not provide much evidence for gene flow due to poor resolution, except for one sub-clade that also contains material of both groups. This could be a signature of a later contact and/or gene flow event. Hence gene flow between populations does not necessarily occur at present; perhaps more likely is that there were further periods of contact at high altitudes during warmer interglacials, and/or low altitudes during glacial maxima.

Adaptive distinctiveness of the three ESUs

Outlier loci identified by BAYESCAN included many that were involved in biosynthetic and metabolic processes of such secondary metabolites as stilbene, coumarin, lignin and L-phenylalanine (Table S7). This might be evidence of adaptive differences among the three lineages, because these compounds, especially stilbene, play key roles in defense mechanisms in plants (Chong, Poutaraud, & Hugueney, 2009). Because of long lifespans, conifers are vulnerable to attack by insects and pathogens, especially the combination of bark beetles and their symbiotic pathogenic fungi (Fettig et al., 2007; Krokene, 2015). Both beetles and fungi can be inhibited by stilbenes, which may be constantly present in bark and/or synthesized following initial attack (Chong et al., 2009; Fettig et al., 2007; Kolosova & Bohlmann, 2012; Krokene, 2015). Although there is little data available for the distribution of insects or fungal pathogens, results from our ongoing analysis of soil microorganisms from *C. chengiana* populations showed an un-even distribution of plant pathogens among the three ESUs (Wang et al., unpublished data). In addition, Stilbenes may also be involved in responses to abiotic stresses like wounding or ozone generated by ultraviolet radiation (Chiron et al., 2000; He, Wu, Pan, & Jiang, 2008; Rosemann, Heller, & Sandermann, 1991). Therefore, evidence detected here of strong selective pressure upon stilbene production mechanisms, implies that each ESU might differ genetically from the others in how it responds to pathogens or abiotic stresses, emphasizing the need to conserve all three to best protect the species.

Interestingly, many of the genes with high PBS_{DDH} were likewise involved in metabolic

and biosynthetic processes of the same compounds, indicating that genes related to these compounds might have played an important role in the adaption of *C. chengiana* to different environments. Comparing to BLJ, DDH is distributed at higher altitudes and might have experienced stronger selection pressures. Those genes identified by PBS as under positive selection in DDH did not include genes with significantly enriched functions directly related to high-altitude adaptation, however many genes that are functionally related to local adaptation did have extreme PBS_{DDH} values and hence might underlie ecological divergence between DDH and BLJ ESU. Of these genes, 34 and 11 were involved in the response to abiotic and biotic stimuli, respectively, and 14 of them were involved in the defense response, all of which might be the signature of DDH ESU to adapt to harsh environment (Table S9). In particular, these genes included six involved in cellular response to DNA damage stimuli, and 13 related to response to radiation (Table S9), which might be of have been important in adapting DDH material to high altitudes, because such habitats are exposed to increased UV radiation which can result in cell and DNA damage (Zeng et al., 2020).

Conservation implications

The three ESUs separated here each face very different conservation issues, so we recommend that each ESU should be assessed independently with regard to its threatened status. Currently, the BLJ ESU has a relatively large extent of occurrence, encompassing Jiuzhaigou County in Sichuan, Wen and Zhouqu County and Wudu district in Gansu. However, its area of occupancy is very narrow, only between 800m to 1500m in the dry valley along Bailongjiang River where there are few steep rocky slopes. These populations are much more accessible to humans than DDH or MJR, and while all areas of *C. chengiana* have suffered from long-term logging (Hao et al., 2006; Zeng & Yang, 1992), the resulting sharp decline in population size may be worst for BLJ, with populations fragmented, and only a few trees remaining. The estimated effective population size by FSC2 is 238,794 (the greatest of the three ESUs), and the Stairway Plot indicates that its effective population size has been stable at approximately 160,000 for the past seven million years (Figure 3a). However, given the long generation times and that most logging is recent, these statistics probably estimate population size before logging. Each habitat of the BLJ ESU should be conserved carefully, and artificial transplanting among fragmented habitats should be undertaken to reduce inbreeding and minimize the bottleneck effect.

Populations in DDH tend to occupy deep slopes near the river, restricting accessibility to loggers. Hence many mature trees of *C. chengiana* remain as part of a wide extent of natural pure forest along the Daduhe River. Noteworthy, our results suggest that a distinct series of locally adapted genetic variations are harbored in DDH populations (Table S8; Figure S5). However, this river is very suitable for hydropower development, with many stations built and others on the way; therefore, large potential habitats of the DDH ESU could be flooded and wiped out (Peng, Li, Wang, Xie, & Cao, 2011). Hence the DDH ESU faces a potentially high risk of extinction, and seed collection for *ex situ* conservation is necessary from populations threatened by development.

Our results suggest that the MJR ESU might be a lineage that experienced an independent evolutionary history following a hybrid origin, making it eligible for protection as a “type 1” taxon (Allendorf et al., 2001). This lineage is now endangered because the extent of suitable

arid habitats is very limited: its range comprises only a small natural forest around Li County plus several fragmented patches in Mao and Wenchuan County. Consistent with this, its current effective population size is the least of the three ESUs as estimated by both FSC2 ($N_2 = 114,433$; Figure 3b) and Stairway Plot (Figure 3a). Our results indicate that this ESU merits conservation in its own right, for which we recommend protection of existing populations and *in situ* augmentation by planting more material. Because each ESU is genetically distinct, any reintroductions or plantings should involve material from the same ESU, to avoid outbreeding depression and preserve genetic distinctness.

From a wider perspective, our study emphasizes the utility of HTS data in conservation genetics for threatened species that have complex genetic structure and evolutionary history. At the same time, our findings also shed light on the formation of lineage diversity in biodiversity hotspots like the HDMs, highlighting the likely roles of hybridization, local adaptation, orogeny and climatic changes.

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Data Accessibility Statement

The transcriptome sequencing data have been deposited in NCBI with the BioProject ID: PRJNA556937 that is publicly accessible at <https://www.ncbi.nlm.nih.gov/bioproject/PRJNA556937>. The filtered SNP matrices file is available in Dryad at <https://doi.org/10.5061/dryad.70rxwdbtf>. The custom scripts have been made available in Supplemental Information (Supplementary Text: Python scripts).

Author Contributions

K.M. and Jianquan Liu designed and supervised this study. Jialiang Li, J.M., W.T., L.Z., D.R. and J.X. managed fieldwork and collected the materials. Jialiang Li and D.R. analyzed the data. Jialiang Li, R.M. and K.M. wrote the manuscript. Jianquan Liu revised the manuscript.

Tables and Figures

Tables

Table 1 Location information of the sampled *Cupressus chengiana* populations (GS = Gansu province, SC = Sichuan province, N = number of individuals analyzed).

Group	Code	N	Location	Latitude (N)	Longitude (E)	Altitude (m)
BLJ	ZR-17	8	Zhouqu, GS	33°52.908'	104°08.156'	1521
	ZR-18	7	Wudu, GS	33°12.030'	105°02.130'	1025
	ZR-20	8	Longnan, GS	33°15.197'	104°59.024'	1634
	ZR-21	7	Wenxian, GS	32°49.364'	104°45.576'	1535
	LXT-18	5	Jiuzhaigou, SC	33°06.946'	104°19.491'	1351
MJR	ZR-01	5	Lixian, SC	31°40.420'	103°49.448'	1500-1938
	ZR-05	7	Lixian, SC	31°23.480'	103°03.680'	2106
	LXT-16	5	Maoxian, CS	31°38.392'	103°48.350'	1742
DDH	ZR-27	7	Jinchuan, SC	31°54.630'	102°01.797'	2410
	ZR-28	6	Maerkang, SC	31°47.460'	101°56.480'	2470
	ZR-30	7	Xiaojin, SC	31°09.790'	102°26.614'	2571
	LXT-05	5	Xiaojin, SC	31°01.749'	102°15.016'	2252
	LXT-08	5	Danba, SC	30°42.954'	101°59.692'	2211

Table 2 Summary of genomic polymorphisms and variants in different *C. chengiana* groups

Parameters		Groups		
		BLJ	MJR	DDH
SNPs	266,884	231,513	228,734	224,658
Private SNPs	-	11,489	3,213	11,913
π	0.0077	0.0069	0.0072	0.0064
Tajima's <i>D</i>	-0.4310	-0.1790	0.0470	-0.1449
		B vs. D	M vs. B	M vs. D
Shared SNPs	-	14,748	27,524	20,245
F_{ST}	-	0.1752	0.1066	0.1397
D_{XY}	-	0.3440	0.3008	0.3104

The π and Tjima's *D* were calculated using the date set without MAF filtering, and the other parameters were calculated based on the data set after MAF control.

Table 3 Inferred demographic parameters for the best-fitting FSC2 model shown in Figure 3b, including 95% confidence intervals.

Parameters	Point estimation	95% confidence intervals	
		Lower bound	Upper bound
N_A	323,866	310,722	330,971
N_1	238,794	227,162	248,078
N_2	114,433	97,828	131,628
N_3	166,952	155,815	172,291
T_1	1,344,300	1,140,450	1,408,600
T_2	4,559,100	4,225,050	4,869,500
α	0.62	0.54	0.65
$M_{1 \leftarrow 2}$	9.22E-06	4.95E-06	1.22E-05
$M_{2 \leftarrow 1}$	3.65E-06	1.37E-06	5.55E-06
$M_{2 \leftarrow 3}$	4.94E-06	3.17E-06	6.44E-06
$M_{3 \leftarrow 2}$	1.00E-05	6.36E-06	1.19E-05
$M_{1 \leftarrow 3}$	9.83E-07	3.29E-07	1.71E-06
$M_{3 \leftarrow 1}$	1.70E-06	6.83E-07	2.49E-06
$MA_{3 \leftarrow 1}$	2.95E-07	1.58E-07	1.43E-06
$MA_{1 \leftarrow 3}$	5.72E-07	1.35E-07	1.35E-06

Parameters included here comprise population size measures (N_A , N_1 , N_2 and N_3 , indicating ancestral population, BLJ, MJR and DDH, respectively), population divergence time (T_2 , years) and hybrid origin time (T_1 , years), hybrid parameter (α), migration per generation after hybridization (M) between each pair of ESUs in each direction, and migration per generation before hybridization (MA) between BLJ and DDH.

959 **Table 4** Results of the ABBA-BABA test. Patterson's *D* value for introgression between
 960 lineages with *Z* score and significance values were shown.

P ₁	P ₂	P ₃	Patterson' <i>D</i>	<i>Z</i> score	<i>p</i> ⁹⁶¹
BLJ	MJR	DDH	0.0780	20.2321	0
DDH	MJR	BLJ	0.1222	28.5418	0

Figure Legends

Figure 1 Geographic distributions of sampled *Cupressus chengiana* populations. Those individuals in the BLJ, MJR and DDH group are distributed in the upper reaches of the Bailongjiang River, the Minjiang River and the Daduhe River, respectively.

Figure 2 Genetic structure and Phylogenetic relationships of the three *C. chengiana* groups (BLJ, MJR, DDH). (a) Admixture proportions of genetic clusters for each individual of the three groups. The scenarios of $K=2$ and $K=3$ are shown, and $K=3$ is the best value according to cross-validation analysis. (b) Principal component analysis (PCA) plot for the 82 *C. chengiana* individuals based on the first two principal components. (c) An ML tree based on 31,527 SNPs in 4DTV of nuclear genome, with three distinct lineages (BLJ, MJR, DDH) detected, among which the relationship between the MJR and BLJ groups is the closest. (d) An ML tree based on 1,251 SNPs of the chloroplast genome, in which the MJR group was not distinguished from the DDH group, while all individuals in the BLJ group form a separate lineage. The supporting values from bootstrap analyses are labeled beside the nodes. Group information is shown in Table 1 and Figure 1.

Figure 3 (a) The detailed population demographic history of BLJ, MJR and DDH over the last 10 million years inferred by Stairway Plot method. Thick lines represent the median, and thin light lines represent the 95% pseudo-CI defined by the 2.5% and 97.5% estimations from the SFS analysis. The periods of the Xixiabangma Glaciation and the Naynayxungla Glaciation are highlighted in gray vertical bars. (b) Maximum likelihood parameter estimates of the best fit models (model3) in FSC2. (c) An ML-bootstrap network for 10,227 orthologous gene trees yielded in PhyloNet with a maximum of two reticulations allowed. The light blue curves represent reticulations with inheritance probabilities behind them. (d) Results of Bayesian outlier analysis for 266,884 SNPs. SNPs with $q\text{-value} < 0.001$ were recognized as outliers. A total of 575 positive outlier SNPs were identified in this analysis.

Figure 4 ENMs for three *C. chengiana* lineages, and identity tests results between paired groups. (a) Current potential distributions of BLJ, MJR and DDH groups, predicted by Maxent. (b) Results of identity tests of three comparisons (BLJ vs. DDH, MJR vs. DDH, MJR vs. BLJ). The grey bars indicate the null distributions of D, while the black bars indicate I. Arrows indicate values of D (gray) and I (black) in actual Maxent runs.